(www.isrec.isb-sib,ch/software/software.html). Identification of surrogates provides tools for partner identification, phenotyping and small molecule discovery. Given that a site directed assay is available at this early stage for the unknown target, high throughout screening allows the rapid identification of reactive small molecules of low target affinity. Combinatorial chemistry, allows for improvements in potency which would then provide small molecules for phenotyping and testing in animal models.

PM

Please amend the following paragraph beginning at page 19, line 34 to page 20,

line 3, to read as follows:

Sequencing of randomly selected clones from the cell library indicated that about 54% of all clones were in-frame. The short FLAG sequence DYKD(SEQ ID NO: 1), was included at the N-terminus as an immunoaffinity tag. In addition, the E-tag epitope (GAPVPYPDPLEPR) (SEQ ID NO: 2) was engineered into the carboxy terminus of the peptide.

Please amend the following paragraph beginning at page 21, line 27, to read as follows:

The panning experiments identified a surrogate peptide, KcB7, with the amino acid sequence RKEMGGGGGPGWSENLFQ(SEQ ID NO: 3). A Blastp search, using several different queries revealed TNFR1 which is the natural biological partner of TNFβ.



Please amend the following section, beginning at page 21, line 34 to page 22, line 28, to

### read as follows:

## BLASTp search results for the TNFB Surrogate peptide KcB7

Query: WSENLFQ(SEQ ID NO: 4)

Database: nr

Sequences producing significant alignments:	Score (bits)	E Value
prf  2102238A tumor necrosis factor alpha inhibitor [Homo s gb AAA36756.1  (M60275) TNF receptor [Homo sapiens pdb 1TNR R Chain R, Tumor Necrosis Factor Receptor P55 pdb 1NCF A Chain A, Binding Protein, Cytokine Mol_id: 1; Mo ref NP_001056.1  tumor necrosis factor receptor 1 (55kD) >g	20 20 20 20 20 20	2419 2419 2419 2419 2419

>prf||2102238A tumor necrosis factor alpha inhibitor [Homo sapiens] Length = 160

Score = 20.4 bits (41), Expect = 2419 Identities = 7/7 (100%), Positives = 7/7 (100%)

Query: 1 WSENLFQ 7(SEQ ID NO: 4)

**WSENLFQ** 

Sbjct: 96 WSENLFQ 102(SEQ ID NO: 4)

>gb|AAA36756.1| (M60275) TNF receptor [Homo sapiens] Length = 453

Score = 20.4 bits (41), Expect = 2419 Identities = 7/7 (100%), Positives = 7/7 (100%)

Query: 1 WSENLFQ 7(SEQ ID NO: 4)

WSENLFQ

Sbjct: 136 WSENLFQ 142(SEQ ID NO: 4)

>pdb|1TNR|R Chain R, Tumor Necrosis Factor Receptor P55 (Extracellular Domain) Complexed With Tumor Necrosis Factor-Beta Length = 139

-3-

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Score = 20.4 bits (41), Expect = 2419 Identities = 7/7 (100%), Positives = 7/7 (100%)

AU

Patternfind search results for the TNF-β Surrogate peptide KcB7

Query sequence:

WSENLFQ(SEQ ID NO: 4)

Please amend the following paragraph beginning at page 22, line 31 to page 23, line 8, to read as follows:

Patternfind search results for the TNF-β Surrogate peptide KcB7

Query sequence:

WSENLFQ(SEQ ID NO: 4)

I. DATABASE:

**NONREDUNDANT** 

Limit

10

gp|M60275|339760|AC886035F969E231 TNF receptor [Homo sapiens]

Occurences: 1

Position: 136 WSENLFQ (SEQ ID NO: 4)

sp|P19438|TNR1\_HUMAN|4CEFBA96D03B8225 (TNFRSF1A..)TUMOR NECROSIS FACTOR RECEPTOR 1 PRECURSOR (TUMOR NECROSIS FACTOR BINDING PROTEIN 1) (TBPI) (P60) (TNF-R1) (TNF-RI) (P55) (CD120A).[Homo sapiens]

Occurences: 1

Position: 136 WSENLFQ(SEQ ID NO: 4)

2 matches found

Please amend the following paragraph beginning at page 23, line 10, to read as follows:

Closer examination of the complementary sequences revealed that the short N-

terminal sequence RKEMG(Piece Of SEQ ID NO: 3) and the C-terminal sequence WSENLFQ

(SEQ ID NO: 4) were identical to regions on TNFR1 (amino acids 77-81 and 107-113

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respectively). These segments corresponded to amino acids within two critical ligand:receptor contact domains. In the case of the N-terminal grouping, the surrogate contained 5 of the 15 amino acids of the 77-81 contact domain whereas in the C-terminal grouping, the surrogate contained 6 of the 9 amino acids identified within the 107-113 contact domain.

Please amend the following section, beginning at page 23, line 20 to page 24, line 5, to read as follows:

## Comparison with human TNFR1 extracellular domain

IYPSGVIGLVPHLGDREKRDSVCPQGKYIHPQNNSICCTKCHKGTYLYNDCPGPGQDTD CRECesgsFTASENHLRhcLscSkCRkeMgQVEISSCTVDRDTVCGCRKNQYRHYWSENLF acFNCSLCLNGTVHLSCQEKQNTVCTCHAGFFLRENECVSCS (SEQ ID NO: 5)



Contact residues are based on Banner et al., (1993) Cell 73: 431-445.

**Bold=** contacted by TNF $\beta$  subunit A **lower case** = contacted by TNF $\beta$  subunit C **italics** = contacted by TNF $\beta$  both subunits A and C Underline = homology to the  $\phi$  clone

TNFβ

LPGVGLTPSAAQTARQHPKMHLAHSTLKPAAHLIGDPSKQNSLLWRANTDRAFLQDGF SLSNNSLLVPTSGIYFVYSQVVFSGKAYSPKAPSspLyLAHEVQLFSsqypfHvPLLSSqKmV YPGLQEPWLHSMYHGAAFQLTQGDQLSThTdGIPHLVLSPSTVFFGAFAL (SEQ ID NO: 6)

**Bold** = TNF $\beta$  subunit A **lower case** = TNF $\beta$  subunit C

Please amend the following section, beginning at page 24, line 8, to read as follows:

### Comparison with human TNFR2 extracellular domain

rKEMGGGGGpgwSENlFQ(SEQ ID NO: 7)



LPAQVAFTPYAPEPGSTCRLREYYDQTAQMCCSKCSPGQHAKVFCTKTSDTVCDSCEDS TYTQLWNWVPECLSCGSRCSSDQVETQACTrEQNRICTCRpgwYCAlSKQEGCRLCAPLR KCRPGFGVARPGTETSDVVCKPCAPGTFSNTTSSTDICRPHQICNVVAIPGNASRDAVCT STSPT(SEQ ID NO: 8)

Please amend the following section beginning at page 26, line 20, to read as follows:

	HCV ALIGNMENTS	
	eIF3:	EDLDNIQTPE-SVLLSAVSGEDTQDRTDRLLLTPWVKFLWESY(SEQ ID NO: 10)
n9	CONSENSUS	TxRLL (SEQ ID NO:11)
171	HCV-NG-D9	T <u>SGE</u> SSG <u>DRT</u> R <u>RVL</u> TSSSART <u>L</u> PN(SEQ ID NO:12)
	HCV-3-F5	$\underline{L}$ VTG $\underline{O}$ F $\underline{P}$ $\underline{S}$ Q $\underline{L}$ LLGGAVCGP—S $\underline{T}$ P $\underline{R}$ L $\underline{T}$ GLCRLSGT(SEQ ID NO:13)
	HCV-3-H8	RRTCGDPAAMLERLSCRAGDYRGASHTGRLLNLRGMHQYP(SEQ ID NO: 14)
	HCV-3-C3	FTTPRHLSGRTVOMMRDSTS(SEQ ID NO: 15)

Please amend the following section beginning at page 26, line 26, to read as follows:

## OUTPUT FROM ADVANCED BLAST SEARCH FOR HCV mRNA SURROGATE QUERY - SEARCH 1:

Query sequence:

TSGESSGDRTRRVLT (SEQ ID NO: 16)

Program:

blastp

Database:

swissprot

Expect value:

10000

Please amend the following section, beginning at page 26, line 33 to page 28, line 35, to read as follows:

#### **OUTPUT**:

Sequences producing significant alignments:	Score E (bits)	Value
sp   P31258   HXAB_CHICK HOMEOBOX PROTEIN HOX-A11 (GHOX-1I) (CH	. 20	477
sp P23116 IF3A MOUSE EUKARYOTIC TRANSLATION INITIATION FACT	. 19	1072
sp   P39690   KHS1 YEAST KILLER TOXIN KHS PRECURSOR (KILLER OF	. 19	1072
sp 083264 NUSG TREPA TRANSCRIPTION ANTITERMINATION PROTEIN	. 19	1072
sp P13079 CARB_STRTH RRNA METHYLTRANSFERASE	19	1072
(CARBOMYCIN-RES		
sp   Q14152   IF3A HUMAN EUKARYOTIC TRANSLATION INITIATION FACT	. 19	1072
sp P39925 AFG3 YEAST MITOCHONDRIAL RESPIRATORY CHAIN COMPLE	. 19	1072
sp P16561 HEMA_VACCT HEMAGGLUTININ PRECURSOR	19	1404
sp P52023 DP3B SYNP7 DNA POLYMERASE III, BETA CHAIN	19	1404
sp   P20978   HEMA_VACCC	19	1404
sp P15989 CA36_CHICK COLLAGEN ALPHA 3 (VI) CHAIN PRECURSOR	19	1404
List truncated here		

>sp|P31258|HXAB\_CHICK HOMEOBOX PROTEIN HOX-A11 (GHOX-11) (CHOX-1.9)

Length = 297

Score = 20.4 bits (41), Expect = 477Identities = 8/11 (72%), Positives = 9/11 (81%)

Query: 2 SGESSGDRTRR 12 (SEQ ID NO: 17)

SG SSG RTR+

Sbjct: 217 SGSSSGQRTRK 227 (SEQ ID NO: 18)

>sp|P23116|IF3A\_MOUSE EUKARYOTIC TRANSLATION INITIATION FACTOR 3 SUBUNIT 10 (EIF-3 THETA)

(EIF3 P167) (EIF3 P180) (EIF3 P185) (P162 PROTEIN)

(CENTROSOMIN) Length = 1344

Score = 19.2 bits (38), Expect = 1072 Identities = 8/13 (61%), Positives = 10/13 (76%)

Query: 2 SGESSGDRTRRVL 14(SEQ ID NO: 19)

SGE + DRT R+L

Sbjct: 133 SGEDTQDRTDRLL 145(SEQ ID NO: 20)

>sp|P39690|KHS1\_YEAST KILLER TOXIN KHS PRECURSOR (KILLER OF HEAT SENSITIVE)

Length = 708

Score = 19.2 bits (38), Expect = 1072Identities = 8/13 (61%), Positives = 10/13 (76%)

Query: 3 GESSGDRTRRVLT 15(SEQ ID NO: 21) G+SSG T+R LT

Sbjct: 98 GKSSGSATKRGLT 110(SEQ ID NO: 22)

>sp|083264|NUSG\_TREPA TRANSCRIPTION ANTITERMINATION PROTEIN NUSG Length = 185

Score = 19.2 bits (38), Expect = 1072Identities = 7/12 (58%), Positives = 9/12 (74%)

Query: 2 SGESSGDRTRRV 13(SEQ ID NO: 23)

+GE GDRT R+

Sbjct: 117 AGEIKGDRTPRI 128(SEQ ID NO: 24)

>sp|P13079|CARB\_STRTH RRNA METHYLTRANSFERASE (CARBOMYCIN-RESISTANCE PROTEIN)

Length = 299

Score = 19.2 bits (38), Expect = 1072 Identities = 8/12 (66%), Positives = 8/12 (66%)

Ouery: 2 SGESSGDRTRRV 13(SEQ ID NO: 23)

SG S DR RRV

Sbjct: 40 SGRSEADRRRRV 51(SEQ ID NO: 25)

>sp|Q14152|IF3A\_HUMAN EUKARYOTIC TRANSLATION INITIATION FACTOR 3 SUBUNIT 10 (EIF-3 THETA)

(EIF3 P167) (EIF3 P180) (EIF3 P185) (KIAA0139) Length = 1382

Score = 19.2 bits (38), Expect = 1072 Identities = 8/13 (61%), Positives = 10/13 (76%)

Query: 2 SGESSGDRTRRVL 14(SEQ ID NO: 19)

SGE + DRT R+L

Sbjct: 133 SGEDTQDRTDRLL 145(SEQ ID NO: 26)

>sp|P39925|AFG3\_YEAST MITOCHONDRIAL RESPIRATORY CHAIN COMPLEXES ASSEMBLY PROTEIN AFG3

(TAT-BINDING HOMOLOG 10) Length = 761

Score = 19.2 bits (38), Expect = 1072Identities = 8/14 (57%), Positives = 10/14 (71%)

Query: 2 SGESSGDRTRRVLT 15(SEQ ID NO: 27)

S +SGD + RVLT

Sbjct: 136 SSNNSGDDSNRVLT 149(SEQ ID NO: 28)

Please amend the following section beginning at page 29, line 1 to page 32, line 30, to read as follows:

OUTPUT FROM ADVANCED BLAST SEARCH FOR HCV mRNA SURROGATE QUERY - SEARCH 2:

**Ouery sequence:** TSGESSGDRTRRVLTSSS(SEQ ID NO: 29)

Program: blastp
Database: swissprot
Expect value: -e 10000

Sequences producing significant alignments:	Score E (bits)	Value
sp Q01728 NAC1_RAT_SODIUM/CALCIUM EXCHANGER 1 PRECURSOR (NA	21	65
spiP70414 NAC1_MOUSE_SODIUM/CALCIUM EXCHANGER 1 PRECURSOR (	21	65
spiP48765 NAC1_BOVIN_SODIUM/CALCIUM EXCHANGER 1 PRECURSOR (	20	190
SDIP48766 NAC1 CAVPO SODIUM/CALCIUM EXCHANGER 1 PRECURSOR (	20	190
SDIP32418INAC1 HUMAN SODIUM/CALCIUM EXCHANGER 1 PRECURSOR (	20	190
spiP23685 NAC1_CANFA_SODIUM/CALCIUM EXCHANGER 1 PRECURSOR (	20	190
sp P48767 NAC1_FELCA_SODIUM/CALCIUM EXCHANGER 1 PRECURSOR (	20	190

```
sp|P08173|ACM4_HUMAN MUSCARINIC ACETYLCHOLINE RECEPTOR M4
                                                              19
                                                                    249
sp|P23116|IF3A_MOUSE_EUKARYOTIC TRANSLATION INITIATION FACT...
                                                                    249
                                                             19
SDIQ14152IIF3A HUMAN EUKARYOTIC TRANSLATION INITIATION FACT...
                                                                    249
                                                              19
sp|P15656|FGF5_MOUSE FIBROBLAST GROWTH FACTOR-5 PRECURSOR (...
                                                             19
                                                                    327
sp|P30042|ES1_HUMAN ES1 PROTEIN HOMOLOG PRECURSOR (PROTEIN ...
                                                              19
                                                                    327
sp|O35491|CLK2_MOUSE_PROTEIN KINASE CLK2
                                                              18
                                                                    428
sp|P49760|CLK2_HUMAN PROTEIN KINASE CLK2
                                                                    428
                                                             18
SPIP15172|MYOD HUMAN MYOBLAST DETERMINATION PROTEIN 1 (MYOG...
                                                             18
                                                                    428
                                                                    428
sp|O75069|Y481_HUMAN HYPOTHETICAL PROTEIN KIAA0481 (HH1480)
                                                             18
sp|P02533|K1CN_HUMAN KERATIN, TYPE I CYTOSKELETAL 14 (CYTOK...
                                                             18
                                                                    561
                                                                    561
spiP30989INTR1 HUMAN NEUROTENSIN RECEPTOR TYPE 1 (NT-R-1) (...
                                                             18
sp|P30551|CCKR_RAT_CHOLECYSTOKININ TYPE A RECEPTOR (CCK-A R...
                                                             18
                                                                    561
spiQ08369|GAT4_MOUSE_TRANSCRIPTION FACTOR GATA-4 (GATA BIND...
                                                                    561
List truncated here...
>sp|Q01728|NAC1 RAT SODIUM/CALCIUM EXCHANGER 1 PRECURSOR
(NA+/CA2+-EXCHANGE PROTEIN 1)
           Length = 971
 Score = 21.2 bits (43), Expect =
 Identities = 9/15 (60%), Positives = 11/15 (73%)
Query: 3
             GESSGDRTRRVLTSS 17(SEQ ID NO: 30)
             GE
                 G RT ++LTSS
Sbjct: 933 GELGGPRTAKLLTSS 947(SEQ ID NO: 31)
>sp|P70414|NAC1 MOUSE SODIUM/CALCIUM EXCHANGER 1 PRECURSOR
(NA+/CA2+-EXCHANGE PROTEIN 1)
           Length = 970
 Score = 21.2 bits (43), Expect =
 Identities = 9/15 (60%), Positives = 11/15 (73%)
Query: 3
             GESSGDRTRRVLTSS 17(SEQ ID NO: 30)
                 G RT ++LTSS
Sbjct: 932 GELGGPRTAKLLTSS 946(SEQ ID NO: 32)
>sp|P48765|NAC1 BOVIN SODIUM/CALCIUM EXCHANGER 1 PRECURSOR
(NA+/CA2+-EXCHANGE PROTEIN 1)
           Length = 970
 Score = 19.6 bits (39), Expect =
 Identities = 8/14 (57%), Positives = 10/14 (71%)
```

```
GESSGDRTRRVLTS 16(SEQ ID NO: 33)
Ouery: 3
           GE G RT ++LTS
Sbjct: 932 GELGGPRTAKLLTS 945(SEQ ID NO: 34)
>sp|P48766|NAC1_CAVPO SODIUM/CALCIUM EXCHANGER 1 PRECURSOR
(NA+/CA2+-EXCHANGE PROTEIN 1)
          Length = 970
 Score = 19.6 bits (39), Expect =
 Identities = 8/14 (57%), Positives = 10/14 (71%)
Query: 3
           GESSGDRTRRVLTS 16(SEQ ID NO: 33)
           GE G RT ++LTS
Sbjct: 932 GELGGPRTAKLLTS 945(SEQ ID NO: 35)
>sp|P32418|NAC1 HUMAN SODIUM/CALCIUM EXCHANGER 1 PRECURSOR
(NA+/CA2+-EXCHANGE PROTEIN 1)
          Length = 970
 Score = 19.6 bits (39), Expect = 190
 Identities = 8/14 (57%), Positives = 10/14 (71%)
Query: 3
           GESSGDRTRRVLTS 16(SEQ ID NO: 33)
           GE G RT ++LTS
Sbjct: 932 GELGGPRTAKLLTS 945(SEQ ID NO: 36)
>sp|P23685|NAC1 CANFA SODIUM/CALCIUM EXCHANGER 1 PRECURSOR
(NA+/CA2+-EXCHANGE PROTEIN 1)
          Length = 970
 Score = 19.6 bits (39), Expect =
 Identities = 8/14 (57%), Positives = 10/14 (71%)
           GESSGDRTRRVLTS 16(SEQ ID NO: 33)
Query: 3
           GE G RT ++LTS
Sbjct: 932 GELGGPRTAKLLTS 945(SEQ ID NO: 37)
>sp|P48767|NAC1_FELCA_SODIUM/CALCIUM_EXCHANGER_1_PRECURSOR
(NA+/CA2+-EXCHANGE PROTEIN 1)
          Length = 970
                                     190
 Score = 19.6 bits (39), Expect =
                                -11-
657408 v1
```

```
GESSGDRTRRVLTS 16(SEQ ID NO: 33)
Query: 3
           GE G RT ++LTS
Sbjct: 932 GELGGPRTAKLLTS 945(SEQ ID NO: 38)
>sp|P08173|ACM4_HUMAN MUSCARINIC ACETYLCHOLINE RECEPTOR M4
          Length = 479
 Score = 19.2 bits (38), Expect =
 Identities = 8/14 (57%), Positives = 13/14 (92%)
Query: 5 SSGDRTRRVLTSSS 18(SEQ ID NO: 39)
          SSG+++ R++TSSS
Sbjct: 10 SSGNQSVRLVTSSS 23(SEQ ID NO: 40)
>sp|P23116|IF3A_MOUSE EUKARYOTIC TRANSLATION INITIATION FACTOR 3
SUBUNIT 10 (EIF-3 THETA)
           (EIF3 P167) (EIF3 P180) (EIF3 P185) (P162 PROTEIN)
           (CENTROSOMIN)
          Length = 1344
 Score = 19.2 bits (38), Expect =
 Identities = 8/13 (61%), Positives = 10/13 (76%)
Query: 2
           SGESSGDRTRRVL 14(SEQ ID NO: 19)
           SGE + DRT R+L
Sbjct: 133 SGEDTQDRTDRLL 145(SEQ ID NO: 20)
>sp|Q14152|IF3A_HUMAN EUKARYOTIC TRANSLATION INITIATION FACTOR 3
SUBUNIT 10 (EIF-3 THETA)
           (EIF3 P167) (EIF3 P180) (EIF3 P185) (KIAA0139)
          Length = 1382
 Score = 19.2 bits (38), Expect = 249
 Identities = 8/13 (61%), Positives = 10/13 (76%)
           SGESSGDRTRRVL 14(SEQ ID NO: 19)
Ouery: 2
           SGE + DRT R+L
Sbjct: 133 SGEDTQDRTDRLL 145(SEQ ID NO: 26)
```

Identities = 8/14 (57%), Positives = 10/14 (71%)

Score = 18.8 bits (37), Expect = 327 Identities = 9/16 (56%), Positives = 10/16 (62%)

Query: 3 GESSGDRTRRVLTSSS 18(SEQ ID NO: 41) G+SSG R R T SS

Sbjct: 39 GDSSGSRGRSSATFSS 54(SEQ ID NO: 42)

>sp|P30042|ES1\_HUMAN ES1 PROTEIN HOMOLOG PRECURSOR (PROTEIN KNP-I) (GT335)

Length = 268

Score = 18.8 bits (37), Expect = 327 Identities = 8/18 (44%), Positives = 11/18 (60%)

Query: 1 TSGESSGDRTRRVLTSSS 18(SEQ ID NO: 43) T G+ S +R VLT S+

Sbjct: 93 TKGQPSEGESRNVLTESA 110(SEQ ID NO: 44)

>sp|035491|CLK2\_MOUSE PROTEIN KINASE CLK2 Length = 499

Score = 18.4 bits (36), Expect = 428 Identities = 8/11 (72%), Positives = 8/11 (72%)

Query: 2 SGESSGDRTRR 12(SEQ ID NO: 17)

S SS DRTRR

Sbjct: 34 SWSSSSDRTRR 44(SEQ ID NO: 45)

>sp|P49760|CLK2\_HUMAN PROTEIN KINASE CLK2 Length = 499

Score = 18.4 bits (36), Expect = 428
Identities = 8/11 (72%), Positives = 8/11 (72%)

Query: 2 SGESSGDRTRR 12(SEQ ID NO: 17)

S SS DRTRR

Sbjct: 34 SWSSSSDRTRR 44(SEQ ID NO: 46)

Please amend the following section beginning at page 33, line 3, to read as follows:

### Output from Patternfind for HCV mRNA surrogate query

Query sequence:

DRTxRLL(SEQ ID NO: 47)

Database:

Nonredundant

Limit:

10

sp|Q14152|IF3A\_HUMAN|485C01B28D67EBBA (EIF3S10)EUKARYOTIC TRANSLATION INITIATION FACTOR 3 SUBUNIT 10 (EIF-3 THETA) (EIF3 P167) (EIF3 P180) (EIF3 P185) (KIAA0139).[Homo sapiens]

Occurrences: 1

Position: 139 DRTDRLL(SEQ ID NO: 48)

sp|P4637 3|FAS1\_RHOFA|A66B6F3DF1286566 (FAS1..)CYTOCHROME P450 FAS1 (EC 1.14.-.-).[Rhodococcus fascians]

Occurrences: 1

Position: 170 DRTARLL(SEQ ID NO: 49)

sp|P23116|IF3A\_MOUSE|F4CAE2169F577712 (EIF3S10..)EUKARYOTIC TRANSLATION INITIATION FACTOR 3 SUBUNIT 10 (EIF-3 THETA) (EIF3 P167) (EIF3 P180) (EIF3 P185) (P162 PROTEIN) (CENTROSOMIN).[Mus musculus]

Occurrences: 1

Position: 139 DRTDRLL(SEQ ID NO: 50)

Please amend the following paragraph, beginning at page 34, line 15, to read as follows:

In addition, several peptides from each pan showed the presence of the **RGG** box, a well-defined RNA-binding motif, as indicated below. **RGG** sequences in each surrogate is in bold and underlined. (SEQ ID NOS 51-65, respectively, in order of appearance)

M1-3-B7	RGLFTEWF <b>RGG</b> SWSNYRVTS
M1-3-E8	TDGGRSVISDNV <u>RGG</u> SRLWLWIRHGSWS <u>Q</u> AWGPQDAWSSK
M1-3-H6	$ ext{RVSSAQPGCTSRVRFRCP}$ $ ext{RGG}$ $ ext{LLFNGVTSTNPKTGLSNAQ}$
M1-4-H1	VVYVGVLSYWPHLSGGGRL <u>O</u> VRCLIG <b>RGG</b> FGC <b>RGG</b>
M2-3-C1	WPPGRTLSDLI <b>RGG</b> AGARGM
M2-3-C9	SSGGLHRWSAL <u>RGG</u> HGHGLA

M2-3-E2	${\tt AMRLKPIAFKGPRAGAGWVEV} \underline{{\tt O}} {\tt PCFAAFRAACT} \underline{{\tt RGG}} {\tt SHHH}$
M2-3-E3	$\verb LHAGWDVTAPRRACKGAO  GPGLHGRFYCH   \textbf{RGG}  LCSGLGRC$
M2-3-E9	${\tt DEQSSLKGKLRGALVRLGMGHAMPH} \underline{{\tt RGG}} {\tt VWPSTGRPSKQG}$
M2-3-H12	${\tt WTPRHGPMRCWRH} \underline{O} {\tt SVFPVGAGPHWALWPIKGP} \underline{\textbf{RGG}} {\tt RTAC}$
M2-NG-C7	${\tt RKTGSNIWLPLYHKVCPASTRAGNG} \underline{{\tt RGG}} {\tt SRFLWGSMQTNC}$
M3-3-B9	${\tt RLQR} \underline{{\tt RGG}} {\tt GAVAVVWVGFGVGLL} \underline{{\tt W}} {\tt GRLLLIILGWVLMWFLS}$
M3-3-C2	QHSEHGGTEWRK <b>RGG</b> MAFAASFLCMRDSYRTTRLRSLLG
M3-3-C7	GTRHVINRVRDSSGVPCKRFGGLQFSQMGKCTIP <b>RGG</b> A
M4-NG-A4	$\verb VL  \frac{\texttt{RGG}}{\texttt{SVGKGSLMWCQEVDWRTGGPRSNLWGLWNGR} \underline{\texttt{O}} \\ \texttt{PPK}$

Please amend the following paragraph beginning at page 34, line 32 to page 35, line 4, to read as follows:

Furthermore, one sequence was found from panning the 20-mer random peptide library on traget M1 that contained the KH motif, which is also a known RNA-binding motifs. The surrogate motif corresponding to the KH domain is in bold and underlined.

```
KH Motif VIGxxGxxF(SEQ ID NO: 66)
M1-3-C6 GVIGGRGLLFPLSGFLHQHR(SEQ ID NO: 67)
```

Please amend the following paragraph beginning at page 37, line 1, to read as follows:

# Sequences of Peptide Binders to Tie-1(SEQ ID NOS 68-72, respectively, in order of appearance)

Consensus:	GxAVVFLDRWGNP
>RPT13	SLWGCSGRAVLFLDSVGNPTGTVRC
>RPT9	RRVD <b>AGGAVVYLDRWGN</b> VSV
>RPT34	VVFLDRWGNPQYLGVKASGG
TT1-G11-R40	GPFSWLFETEWGNPKTVPFGADRWNRHGRWDPGPVSDYGT

Please amend the following paragraph beginning at page 38, line 33 to page 39, line 2 to read as follows:

The complete ORF of the Tie-1 gene is cloned from fetal human brain (Clontech Quick-Clone cDNA) or fetal human heart using the following primers:

- 5' Tie-1 forward: GGT CGG CCT CTG GAG TAT GGT CTG(SEQ ID NO: 76)
- 3' Tie-1 reverse: TCC TTG AGG CAG CTT AAG TCA GAG(SEQ ID NO: 77)

Please amend the following paragraph beginning at page 39, line 3, to read as follows:

The complete ORF of the EGFR gene is cloned from the above libraries or from a placental cDNA library (Clontech Placenta Marathon ready cDNA) using the following primers:

- 5' EGFR forward: GGA GCA GCG ATG CGA CCC TC(SEQ ID NO: 78)
- 3' EGFR reverse: GGT CCT GGG TAT CGA AAG AGT CTG G(SEQ ID NO:

79)

Please amend the following paragraph, beginning at page 39, line 10, to read as follows:

In the chimeric receptor, the extracellular and transmembrane regions of Tie-1 are joined to the cytoplasmic kinase domain of the EGFR with an NHE I site which will add the amino acids alanine and serine at the junction. The primers for generating the chimeric receptor are the following primers (with the NHE site underlined):

EGFR forward: GCG CTG CTA GCC GAA GGC GCC ACA TCG TTC(SEQ ID

NO: 80)

Tie-1 reverse: GCT GCT GCT AGC GAT GCA CAC CAG GGT TAA AAG G

(SEQ ID NO: 81)

Should any fees be required by the filing of this Response, Amendment, Statement, and Sequence Listing, authorization is hereby given to charge the amount of any such fee(s) that is/are properly assessable in this application to Deposit Account No. 13-4500, Docket No. 2598-4004US1. A DUPLICATE COPY OF THIS SHEET IS ATTACHED.

Respectfully submitted,

MORGAN & FINNEGAN, L.L.P.

Dated: November 30, 2001

Kenneth H. Sonnenfeld

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